Amendments to the Claims

Following is a complete set of claims as amended with this Response. This complete set of claims excludes cancelled claim 35 and includes amended claims 25-34.

1. (Withdrawn) A method of performing electrophysiological testing in a cardiac stimulation device capable of delivering non-invasive programmed stimulation, comprising:

detecting a cardiac event in a cardiac chamber;

implementing an electrophysiological testing scheme upon detection of the cardiac event occurring in the cardiac chamber; and

delivering a predetermined sequence of stimulation pulses to the cardiac chamber as dictated by the testing scheme.

- 2. (Withdrawn) The method of claim 1, wherein implementing the testing scheme is performed during a refractory period that follows the detected cardiac event.
- 3. (Withdrawn) The method of claim 2, wherein implementing the testing scheme includes switching from a standard operating mode to a non-invasive programmed stimulation mode.
- 4. (Withdrawn) The method of claim 3, further including receiving an external command that triggers the onset of the non-invasive programmed stimulation.
- 5. (Withdrawn) The method of claim 3, wherein detecting the cardiac event includes detecting an intrinsic event in the cardiac chamber being tested.
- 6. (Withdrawn) The method of claim 5, wherein detecting an intrinsic event includes detecting an intrinsic depolarization occurring in one of an atrial cardiac chamber and a ventricular cardiac chamber.

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- 7. (Withdrawn) The method of claim 3, wherein detecting the cardiac event includes detecting a stimulated event in the cardiac chamber being tested.
- 8. (Withdrawn) The method of claim 7, wherein detecting a stimulated event includes detecting one of an atrial stimulation pulse and a ventricular stimulation pulse.
- 9. (Withdrawn) The method of claim 3, further including providing a recovery delay following the non-invasive programmed stimulation.
- 10. (Withdrawn) The method of claim 9, further comprising starting a second refractory period following the expiration of the recovery delay if no intrinsic event is detected during the recovery delay.
- 11. (Withdrawn) The method of claim 10, further including effecting a transfer from the non-invasive programmed stimulation mode to the standard operating mode during the second refractory period.
- 12. (Withdrawn) The method of claim 1, further including blanking sensing circuitry of non-tested cardiac chambers during the delivery of the sequence of stimulation pulses in the cardiac chamber being tested.
- 13. (Withdrawn) The method of claim 1, further including providing back-up ventricular stimulation whenever atrial non-invasive programmed stimulation is performed; and

wherein providing back-up ventricular stimulation includes providing backup ventricular stimulation at a programmed rate that is decoupled from the atrial non-invasive programmed stimulation.



- 14. (Withdrawn) The method of claim 9, further comprising starting a refractory period if an intrinsic event is sensed in the recovery period.
- 15. (Original) A stimulation device capable of performing electrophysiological testing by delivering non-invasive programmed stimulation, comprising:

a discriminator that senses a cardiac event in a cardiac chamber being tested:

timing circuitry, coupled to the discriminator, that triggers an onset of the non-invasive programmed stimulation based on a detected cardiac event occurring in the cardiac chamber being tested;

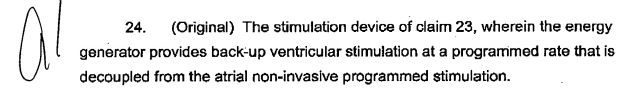
a controller, connected to the timing circuitry that executes a transfer between a first and a second stimulation mode; and

an energy generator connected to the discriminator, the timing circuitry and the controller, the generator is controlled by the controller to deliver a sequence of stimulation pulses to the cardiac chamber being tested in response to the detected cardiac event.

- 16. (Original) The stimulation device of claim 15, wherein the timing circuitry sets a refractory period that follows a triggering detected cardiac event; and wherein the controller executes the transfer during the refractory period.
- 17. (Original) The stimulation device of claim 16, wherein the controller executes the transfer between the first and the second stimulation mode by switching from a standard operating mode to a non-invasive programmed stimulation mode.
- 18. (Original) The stimulation device of claim 17, further including a programmer that generates an external command; and

wherein the timing circuitry triggers the onset of the non-invasive programmed stimulation in response to the external command.

- 19. (Original) The stimulation device of claim 17, wherein the discriminator detects any one of an atrial intrinsic event, ventricular intrinsic event, an atrial stimulated event, or a ventricular stimulated event in the cardiac chamber being tested.
- 20. (Original) The stimulation device of claim 17, wherein the timing circuitry further sets a recovery delay at the expiration of the non-invasive programmed stimulation.
- 21. (Original) The stimulation device of claim 20, wherein the timing circuitry is operative to start a second refractory period following the expiration of the recovery delay if no intrinsic event is detected during the recovery delay.
- 22. (Original) The stimulation device of claim 21, wherein the controller further effects a transfer from the non-invasive programmed stimulation mode to the standard operating mode during the second refractory period.
- 23. (Original) The stimulation device of claim 15, wherein the energy generator further provides back-up ventricular stimulation whenever atrial non-invasive programmed stimulation is performed.



25. (Currently Amended) A stimulation device capable of performing electrophysiological testing by delivering non-invasive programmed stimulation, comprising:

means for detecting sensing circuitry to detect a cardiac event in a cardiac chamber to be tested:

means for implementing a controller coupled to the sensing circuitry, the controller to implement an electrophysiological testing scheme in response to detection of the cardiac event; and

means for delivering a pulse generator coupled to the controller, the pulse generator to deliver a sequence of stimulation pulses to the cardiac chamber as dictated by the testing scheme.

26. (Currently Amended) The stimulation device of claim 25, further comprising means for setting a refractory period that follows the detected cardiac event wherein the controller comprises a timing control circuitry coupled to the sensing circuitry, the timing control circuitry to trigger an onset of the non-invasive programmed stimulation based on the detected cardiac event occurring in the cardiac chamber being tested; and

wherein the implementing-means controller implements the testing scheme during the <u>a</u> refractory period.

27. (Currently Amended) The stimulation device of claim 26, further comprising means for switching wherein the electrophysiological testing scheme comprises a transfer from a standard operating mode to a non-invasive programmed stimulation mode.

28. (Currently Amended) The stimulation device of claim 27, further including means for detecting cardiac events in the chamber being tested; and

wherein the detecting means senses sensing circuitry detects any one of an atrial intrinsic event, ventricular intrinsic event, an atrial stimulated event, or a ventricular stimulated event in the cardiac chamber being tested.

- 29. (Currently Amended) The stimulation device of claim 27, further comprising means for setting wherein the timing control circuitry further sets a recovery delay at the expiration of the non-invasive programmed stimulation.
- 30. (Currently Amended) The stimulation device of claim 29, further comprising means for starting wherein the timing control circuitry is operative to start a second refractory period following the expiration of the recovery delay if no intrinsic event is sensed during the recovery delay.
- 31. (Currently Amended) The stimulation device of claim 30, further comprising means for effecting wherein the controller further effects a transfer from the non-invasive programmed stimulation mode to the standard operating mode during the second refractory period.
- 32. (Currently Amended) The stimulation device of claim 25, wherein the delivering means pulse generator further provides back-up ventricular stimulation whenever atrial non-invasive programmed stimulation is performed.
- 33. (Currently Amended) The stimulation device of claim 32, wherein the delivering means pulse generator provides back-up ventricular stimulation at a programmed rate that is decoupled from the atrial non-invasive programmed stimulation.

- 34. (Currently Amended) The stimulation device of claim 25, further including means for effecting wherein the controller further effects a transfer from the test mode to a normal mode if a failure occurs during the non-invasive programmed stimulation.
 - 35. (Cancelled)

